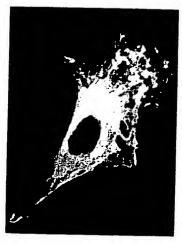
MOLECULAR CELL BIOLOGY

JAMES DARNELL

Vincent Astor Professor Rockefeller University

HARVEY LODISH

Sember of the Whitehead Institute for Biomedical Research Professor of Biology, Massachusetts Institute of Technology



DAVID BALTIMORE

Director of the Whitehead Institute for Biomedical Research Professor of Biology, Massachusetts Institute of Technology

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making a total of eight types of immunoglobulin. Attached to all the H chains are asparagine-linked carbohydrate chains.

Every individual antibody molecule has one type of L chain and one type of H chain. The chains are held together by disulfide bonds to form a monomer, and two monomers are linked by disulfide bonds to form the basic dimeric structure of the molecule (Figure 24-15). Within each chain, units made of about 110 amino acids fold up to form compact domains. Each domain has a single internal disulfide bond which holds it together. An L chain has two domains, H chains have four or five domains. The first two N-terminal domains of the H chains interact with the two L-chain domains, producing a compact unit that acts as the binding region of the antibody. In most H chains, a hinge region consisting of a small number of

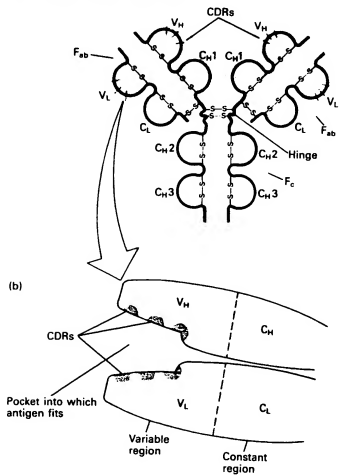
(a)

V_H = Variable region heavy chain

CH = Constant region heavy chain

V_L = Variable region light chain

C_L = Constant region heavy chain



amino acids is found after the first two domains. The hinge is flexible and allows the binding regions to move freely relative to the rest of the molecule. At the hinge region are located the cysteine residues whose SH group are linked to form the —S—S— bridges between the two monomer units of the antibody dimer. The hinge region are the places most susceptible on the molecule to the action of protease; light protease treatment can split an antibody into two pieces, called F_{ab} and F_c fragment. The F_{ab} portion has the antigen-binding site, the F_c portion has the effector regions (see Figure 24-15).

The very first amino acid sequences determined on chains from human myelomas made it clear that N-terminal domain has a very variable structure withe C-terminal domain has a quite constant structure. I N-terminal domain is called the variable region and C-terminal domain is called the constant region (1) 24-15). H-chain sequences show the same division within each class the N-terminal domain is highly vible and the C-terminal domains have a constant quence. The variable domains of L and H chambound to one another. In fact, they interact close! form a single compact unit (Figure 24-15). This unit antibody-binding site, the region of the antibody is cule that binds to antigen. This can be demonstrated the use of an antigen that contains a reactive chogroup: the reactive group on the antigen will form lent bond to the variable domain of the H or 1. showing that the variable domains form the arr binding pocket.

Domain structure and the comple mentarity-determining regions (CDRs) of anni body. (a) The molecule is organized in disulfide bonded 110 amino acid domains, four in the heavy chain and two in the light chain. The far thest N-terminal domain of each chain is variable in sequence (V region): the other domains are constant in sequence (regions C_H1 , C_H2 , and $C_{H^{3/111}}$ the heavy chain and C_L in the light chain). When the molecule is protease-digested, cutting of the hinge region connecting CH1 and CH2, the most sensitive spot on the molecule, splits the molecule into two parts, Fab (the antigen-binding dom.un and Fc (the effector region). Within the V region are segments of highly variable sequence constitution ing the CDRs, the amino acids that actually contact the antigen. (b) Antibody binding site, the if coordinate effort of the CDRs from light and heavy chains, a binding pocket is formed into which a specific antigen fits.

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